

Patient Name	Affix barcode label of Patient's sample here
Date of Birth	

## Form instructions:

- ✓ Specify the testing to be performed on pages 1-2 and sign at the bottom of page 2
- ✓ Provide patient's clinical information on page 3
- ✓ Provide patient demographics on page 4

Comprehensive Analyses	
<input type="radio"/> Genomic Unity® Whole Genome Analysis (CP001) <input type="radio"/> Singleton <input type="radio"/> Duo <input type="radio"/> Trio	Provides whole genome sequence analysis of exonic regions, intronic regions and regulatory variants, genome wide copy number changes, deletions, duplications, inversions, regions of homozygosity and mobile element insertions of the nuclear genome, sequence with heteroplasmy, and deletion analyses of the mitochondrial genome, standard tandem repeat expansion analysis of 24 genes, with an option to add <i>HTT</i> and <i>JPH3</i> with special consent. See full test information: <a href="https://www.variantyx.com/whole-genome-analysis">https://www.variantyx.com/whole-genome-analysis</a> .
<input type="radio"/> Genomic Unity® Exome Plus Analysis (CP010) <input type="radio"/> Singleton <input type="radio"/> Duo <input type="radio"/> Trio	Provides whole exome sequence analysis of exonic regions, characterized intronic regions and regulatory variants, genome wide copy number changes, deletions, duplications, inversions, regions of homozygosity and mobile element insertions of the nuclear genome, sequence with heteroplasmy, and deletion analyses of the mitochondrial genome, standard tandem repeat expansion analysis of 24 genes, with an option to add <i>HTT</i> and <i>JPH3</i> with special consent. See full test information: <a href="https://www.variantyx.com/solutions/diagnostic-testing/genomic-unity-analyses/exome-plus-analysis/">https://www.variantyx.com/solutions/diagnostic-testing/genomic-unity-analyses/exome-plus-analysis/</a>
Other Comprehensive Analyses	
<input type="radio"/> Genomic Unity® Exome Analysis (CP002) <input type="radio"/> Singleton <input type="radio"/> Duo <input type="radio"/> Trio <i>Optional add-on tests</i> <input type="radio"/> Genomic Unity® Mitochondrial Genome Analysis (CP003) <input type="radio"/> Genomic Unity® Constitutional Genome-Wide Copy Number Variant Analysis (CP004) <input type="radio"/> If all 3 analyses (CP002, CP003, CP004) are selected please provide a Unified Whole Genome Analysis report.	Provides whole exome sequence analysis of exonic regions, characterized intronic regions and regulatory variants, including tandem repeat expansion analysis of 24 genes, with an option to add <i>HTT</i> and <i>JPH3</i> with special consent. See full test information: <a href="https://www.variantyx.com/exome-analysis">https://www.variantyx.com/exome-analysis</a> .
<input type="radio"/> Genomic Unity® Mitochondrial Genome Analysis (CP003)	Provides sequence analysis with heteroplasmy and deletion analysis of the mitochondrial genome. See full test information: <a href="https://www.variantyx.com/mito-genome-analysis">https://www.variantyx.com/mito-genome-analysis</a> .
<input type="radio"/> Genomic Unity® Constitutional Genome-Wide Copy Number Variant Analysis (CP004) <i>Optional add-on variant detection (select below)</i> <input type="radio"/> Single nucleotide variants and small deletion/insertions (<50 bp) that intersect reportable copy number variants.	Provides constitutional genome-wide copy number variant analysis, deletions, duplications, inversions, regions of homozygosity, and mobile element insertions of the nuclear genome. See full test information: <a href="https://www.variantyx.com/cnv-analysis">https://www.variantyx.com/cnv-analysis</a> .
Phenotype Based Comprehensive Analyses	
<input type="radio"/> Genomic Unity® Comprehensive Mitochondrial Disorders Analysis (MD001)	Provides sequence analysis with heteroplasmy and deletion analysis of the mitochondrial genome and sequence and duplication/deletion analysis of 335 nuclear genes related to mitochondrial disorders. See full test information: <a href="https://www.variantyx.com/solutions/diagnostic-testing/genomic-unity-analyses/mitochondrial-analysis/">https://www.variantyx.com/solutions/diagnostic-testing/genomic-unity-analyses/mitochondrial-analysis/</a>
<input type="radio"/> Genomic Unity® Intellectual Disability Analysis (NR001)	Provides genome-wide copy number variant analysis, tandem repeat expansion analysis of <i>FMR1</i> and <i>AFF2</i> , full gene sequence and duplication/deletion analysis of genes related to intellectual disability: <i>ADNP</i> , <i>CHD2</i> , <i>FOXP1</i> , <i>FOXP2</i> , <i>GRIN2A</i> , <i>GRIN2B</i> , <i>NLGN3</i> , <i>NLGN4X</i> , <i>MECP2</i> , <i>SHANK3</i> and <i>PTEN</i> . See full test information: <a href="https://www.variantyx.com/intellectual-disability-analysis">https://www.variantyx.com/intellectual-disability-analysis</a> .
<input type="radio"/> Genomic Unity® Comprehensive Ataxia Analysis (NR002)	Provides sequence and duplication/deletion analysis of 51 genes related to ataxia as well as tandem repeat expansion analysis of <i>ATN1</i> , <i>ATXN1</i> , <i>ATXN2</i> , <i>ATXN3</i> , <i>ATXN7</i> , <i>ATXN8OS</i> , <i>ATXN10</i> , <i>CACNA1A</i> , <i>FXN</i> , <i>NOP56</i> , <i>PPP2R2B</i> , <i>TBP</i> . See full test information: <a href="https://www.variantyx.com/ataxia-analysis">https://www.variantyx.com/ataxia-analysis</a> .
<input type="radio"/> Genomic Unity® Ataxia Repeat Expansion Analysis (NR003)	Provides sequence, duplication/deletion analysis and tandem repeat expansion analysis of ataxia-associated genes <i>ATN1</i> , <i>ATXN1</i> , <i>ATXN2</i> , <i>ATXN3</i> , <i>ATXN7</i> , <i>ATXN8OS</i> , <i>ATXN10</i> , <i>CACNA1A</i> , <i>FXN</i> , <i>NOP56</i> , <i>PPP2R2B</i> , <i>TBP</i> . See full test information: <a href="https://www.variantyx.com/ataxia-repeat-analysis">https://www.variantyx.com/ataxia-repeat-analysis</a> .
<input type="radio"/> Genomic Unity® Epilepsy Analysis (NR004)	Provides sequence and duplication/deletion analysis of genes related to seizures as well as tandem repeat expansion analysis of <i>AFF2</i> , <i>CSTB</i> , <i>DIP2B</i> , <i>FMR1</i> . See full test information: <a href="https://www.variantyx.com/epilepsy-analysis">https://www.variantyx.com/epilepsy-analysis</a> .
<input type="radio"/> Genomic Unity® Motor Neuron Disorders Analysis (NR005)	Provides sequence and duplication/deletion analysis of genes related to motor neuron disorders as well as tandem repeat expansion analysis of <i>AR</i> , <i>C9ORF72</i> . See full test information: <a href="https://www.variantyx.com/motor-neuron-analysis">https://www.variantyx.com/motor-neuron-analysis</a> .



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### Phenotype Based Comprehensive Analyses (cont.)

- |   |  |
|---|--|
| <input type="radio"/> Genomic Unity® Movement Disorders Analysis (NR006)      | Provides sequence and duplication/deletion analysis of genes related to movement disorders as well as tandem repeat expansion analysis of <i>ATN1</i> , <i>ATXN1</i> , <i>ATXN2</i> , <i>ATXN3</i> , <i>ATXN7</i> , <i>ATXN8OS</i> , <i>ATXN10</i> , <i>C9ORF72</i> , <i>CACNA1A</i> , <i>FXN</i> , <i>NOP56</i> , <i>NOTCH2NLC</i> , <i>PPP2R2B</i> , <i>TBP</i> . Option to add on tandem repeat expansion of <i>HTT</i> and <i>JPH3</i> with special consent.<br>See full test information: <a href="https://www.variantyx.com/movement-analysis">https://www.variantyx.com/movement-analysis</a> . |
| <input type="radio"/> Genomic Unity® Neuromuscular Disorders Analysis (NR007) | Provides sequence and duplication/deletion analysis of genes related to neuromuscular disorders as well as tandem repeat expansion analysis of the <i>CNBP</i> and <i>DMPK</i> genes.<br>See full test information: <a href="https://www.variantyx.com/neuromuscular-analysis">https://www.variantyx.com/neuromuscular-analysis</a> .  |
| <input type="radio"/> Genomic Unity® Muscular Dystrophy Analysis (NR008)      | Provides sequence and duplication/deletion analysis of genes related to muscular dystrophies.<br>See full test information: <a href="https://www.variantyx.com/md-analysis">https://www.variantyx.com/md-analysis</a> .  |
| <input type="radio"/> Genomic Unity® Neuropathies Analysis (NR009)            | Provides sequence and duplication/deletion analysis of genes related to neuropathies.<br>See full test information: <a href="https://www.variantyx.com/neuropathies-analysis">https://www.variantyx.com/neuropathies-analysis</a> .  |

### Custom Analysis Select when you want to specify the genes analyzed

- |  |   |
|--|---|
| <input type="radio"/> Genomic Unity® Custom Analysis (CA001) | Provides results that are filtered from Genomic Unity® Whole Genome Analysis. Test results include sequence, duplication/deletion analysis and tandem repeat analysis (when relevant) for the specific genes requested.<br>See the list of genes available for this analysis: <a href="https://www.variantyx.com/custom-analysis">https://www.variantyx.com/custom-analysis</a> . |
|--|---|

List the gene(s) to be included in the analysis. If more room is required, please attach a separate page:

The selected genes included in this custom analysis are filtered from a whole genome backbone whereby variants outside the regions of interest are masked, therefore the performance characteristics are based on Genomic Unity® Whole Genome Analysis. The selected genes may: (1) have not been curated and assessed for clinical relevance and utility; (2) have not been sequenced completely (not fully covered) and therefore pathogenic variants in uncovered regions may not be identified; (3) have variants that are not identified or identified with reduced confidence by the Variantyx platform, included but not limited to non-unique genomic regions and high population frequency variants; and/or (4) have variants that require special interpretation that may not be reported.

### Other Analyses Select from additional analyses offered online at [Genomic Unity® Analyses](#)

Test code:

Test name:

### Optional Reflex

In case the targeted analysis selected does not yield a positive result, please reflex to Genomic Unity® comprehensive analyses (select all that apply).  
See full test information: [Genomic Unity® Analyses](#)

- ☐ Genomic Unity® Exome Analysis (CP002)  
☐ Genomic Unity® Mitochondrial Genome Analysis (CP003)  
☐ Genomic Unity® Constitutional Genome-Wide Copy Number Variant Analysis (CP004)

☐ If all 3 analyses (CP002, CP003, CP004) are selected please provide a Unified Whole Genome Analysis report.

☐ Singleton ☐ Duo ☐ Trio

### Healthcare Provider's Statement

By my signature below, I attest that I am the referring physician, an authorized healthcare provider for the patient, or procurator thereof and this testing is medically necessary for diagnosis and/or treatment of the patient. I attest that the patient (or guardian) has been appropriately consented about the test including possible results and outcomes, has been given the opportunity to ask questions about the testing and/or seek genetic counseling, and agrees to allow an independent genetic counselor facilitated through a third party, DNAVisit, to provide pre-test and/or post-test genetic counseling, if required by the insurer and/or referring institution. I attest that the patient (or guardian) has voluntarily consented to testing performed by Variantyx for diagnostic purposes through both oral and written consent.

Healthcare provider signature \_\_\_\_\_

Date \_\_\_\_\_



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Please specify the patient's phenotypes using 1° to indicate the most important primary phenotypes and 2° to indicate less important secondary phenotypes. ICD-10 codes must be specified here and/or in attached clinical notes.

### Clinical Information

ICD-10 Code(s)\*

### Patient Phenotypes

	1° 2°	Phenotype	Age of onset		1° 2°	Phenotype	Age of onset			
Development/Behavior	<input type="radio"/>	<input type="radio"/>	Developmental regression		Constitutional	<input type="radio"/>	<input type="radio"/>	Cleft lip		
	<input type="radio"/>	<input type="radio"/>	Global developmental delay			<input type="radio"/>	<input type="radio"/>	Cleft palate		
	<input type="radio"/>	<input type="radio"/>	Intellectual disability			<input type="radio"/>	<input type="radio"/>	Syndactyly		
	<input type="radio"/>	<input type="radio"/>	Delayed fine motor development			<input type="radio"/>	<input type="radio"/>	Polydactyly		
	<input type="radio"/>	<input type="radio"/>	Delayed gross motor development			<input type="radio"/>	<input type="radio"/>	Failure to thrive		
	<input type="radio"/>	<input type="radio"/>	Delayed speech and language development			<input type="radio"/>	<input type="radio"/>	Macrocephaly		
	<input type="radio"/>	<input type="radio"/>	Speech articulation difficulties			<input type="radio"/>	<input type="radio"/>	Microcephaly		
	<input type="radio"/>	<input type="radio"/>	Autism spectrum disorder			<input type="radio"/>	<input type="radio"/>	Obesity		
	<input type="radio"/>	<input type="radio"/>	Self-injurious behavior			<input type="radio"/>	<input type="radio"/>	Short stature		
	<input type="radio"/>	<input type="radio"/>	Stereotypy			<input type="radio"/>	<input type="radio"/>	Tall stature		
Brain Anomalies	<input type="radio"/>	<input type="radio"/>	Brain atrophy		Ophthalmology/Auditory	<input type="radio"/>	<input type="radio"/>	Blindness		
	<input type="radio"/>	<input type="radio"/>	Cerebellar hypoplasia			<input type="radio"/>	<input type="radio"/>	Cataracts		
	<input type="radio"/>	<input type="radio"/>	Cortical dysplasia			<input type="radio"/>	<input type="radio"/>	Coloboma		
	<input type="radio"/>	<input type="radio"/>	Encephalocele			<input type="radio"/>	<input type="radio"/>	External ophthalmoplegia		
	<input type="radio"/>	<input type="radio"/>	Holoprosencephaly			<input type="radio"/>	<input type="radio"/>	Optic atrophy		
	<input type="radio"/>	<input type="radio"/>	Hydrocephalus			<input type="radio"/>	<input type="radio"/>	Ptosis		
	<input type="radio"/>	<input type="radio"/>	Lissencephaly			<input type="radio"/>	<input type="radio"/>	Rod-cone dystrophy		
	<input type="radio"/>	<input type="radio"/>	Molar tooth sign			<input type="radio"/>	<input type="radio"/>	Visual impairment		
	<input type="radio"/>	<input type="radio"/>	Periventricular leukomalacia			<input type="radio"/>	<input type="radio"/>	Aminoglycoside-induced hearing loss		
	<input type="radio"/>	<input type="radio"/>	Polymicrogyria			<input type="radio"/>	<input type="radio"/>	External ear malformation		
Neurological	<input type="radio"/>	<input type="radio"/>	Abnormal nerve conduction velocity		Cardiac	<input type="radio"/>	<input type="radio"/>	Arrhythmia		
	<input type="radio"/>	<input type="radio"/>	Ataxia			<input type="radio"/>	<input type="radio"/>	Cardiomyopathy		
	<input type="radio"/>	<input type="radio"/>	Spasticity			<input type="radio"/>	<input type="radio"/>	Syncope		
	<input type="radio"/>	<input type="radio"/>	Chorea			<input type="radio"/>	<input type="radio"/>	Tetralogy of Fallot		
	<input type="radio"/>	<input type="radio"/>	Dystonia		Gastrointestinal	<input type="radio"/>	<input type="radio"/>	Aganglionic megacolon		
	<input type="radio"/>	<input type="radio"/>	Foot dorsiflexor weakness			<input type="radio"/>	<input type="radio"/>	Constipation		
	<input type="radio"/>	<input type="radio"/>	Headache			<input type="radio"/>	<input type="radio"/>	Diarrhea		
	<input type="radio"/>	<input type="radio"/>	Neurodegeneration			<input type="radio"/>	<input type="radio"/>	Elevated hepatic transaminases		
	<input type="radio"/>	<input type="radio"/>	Motor axonal neuropathy			<input type="radio"/>	<input type="radio"/>	Gastroesophageal reflux		
	<input type="radio"/>	<input type="radio"/>	Pes cavus			<input type="radio"/>	<input type="radio"/>	Gastroschisis		
<input type="radio"/>	<input type="radio"/>	Reduced deep tendon reflexes		<input type="radio"/>	<input type="radio"/>	Omphalocele				
<input type="radio"/>	<input type="radio"/>	Seizures		<input type="radio"/>	<input type="radio"/>	Pyloric stenosis				
<input type="radio"/>	<input type="radio"/>	Sleep apnea		<input type="radio"/>	<input type="radio"/>	Tracheoesophageal fistula				
<input type="radio"/>	<input type="radio"/>	Stroke-like episodes		<input type="radio"/>	<input type="radio"/>	Vomiting				
<input type="radio"/>	<input type="radio"/>	Tremor		Genitourinary	<input type="radio"/>	<input type="radio"/>	Abnormal renal morphology			
<input type="radio"/>	<input type="radio"/>	Vocal cord paresis			<input type="radio"/>	<input type="radio"/>	Ambiguous genitalia			
Muscular	<input type="radio"/>	<input type="radio"/>	Dysphagia			<input type="radio"/>	<input type="radio"/>	Cryptorchidism		
	<input type="radio"/>	<input type="radio"/>	Exercise intolerance			<input type="radio"/>	<input type="radio"/>	Hydronephrosis		
	<input type="radio"/>	<input type="radio"/>	Hypertonia			<input type="radio"/>	<input type="radio"/>	Hypospadias		
	<input type="radio"/>	<input type="radio"/>	Hypotonia			<input type="radio"/>	<input type="radio"/>	Renal agenesis		
	<input type="radio"/>	<input type="radio"/>	Muscle fasciculations		Skeletal	<input type="radio"/>	<input type="radio"/>	Abnormal vertebral morphology		
	<input type="radio"/>	<input type="radio"/>	Muscle wasting			<input type="radio"/>	<input type="radio"/>	Clubfoot		
<input type="radio"/>	<input type="radio"/>	Muscle weakness		<input type="radio"/>		<input type="radio"/>	Craniosynostosis			
<input type="radio"/>	<input type="radio"/>	Muscular dystrophy		<input type="radio"/>	<input type="radio"/>	Multiple joint contractures				
<input type="radio"/>	<input type="radio"/>	Myotonia		<input type="radio"/>	<input type="radio"/>	Scoliosis				
Metabolic	<input type="radio"/>	<input type="radio"/>	Aciduria		Skin	<input type="radio"/>	<input type="radio"/>	Abnormality of connective tissue		
	<input type="radio"/>	<input type="radio"/>	Abnormal CPK circulation concentration			<input type="radio"/>	<input type="radio"/>	Abnormality of skin pigmentation		
	<input type="radio"/>	<input type="radio"/>	Decreased plasma carnitine			<input type="radio"/>	<input type="radio"/>	Abnormality of temperature regulation		
	<input type="radio"/>	<input type="radio"/>	Elevated serum alanine aminotransferase			<input type="radio"/>	<input type="radio"/>	Ichthyosis		
	<input type="radio"/>	<input type="radio"/>	Increased serum pyruvate			Other phenotypes				
	<input type="radio"/>	<input type="radio"/>	Ketosis							
<input type="radio"/>	<input type="radio"/>	Lactic acidosis								
<input type="radio"/>	<input type="radio"/>									
Endocrine	<input type="radio"/>	<input type="radio"/>	Adrenal hyperplasia							
	<input type="radio"/>	<input type="radio"/>	Adrenal insufficiency							
	<input type="radio"/>	<input type="radio"/>	Cushing syndrome							
	<input type="radio"/>	<input type="radio"/>	Diabetes Mellitus Type I							
	<input type="radio"/>	<input type="radio"/>	Diabetes Mellitus Type II							
	<input type="radio"/>	<input type="radio"/>	Hypothyroidism							
	<input type="radio"/>	<input type="radio"/>	Hypoparathyroidism							
	<input type="radio"/>	<input type="radio"/>	Hypogonadism							
	<input type="radio"/>	<input type="radio"/>	Paraganglioma							





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Patient Information					
First Name	Last Name		MI	DOB	Genetic Sex <input type="radio"/> Male <input type="radio"/> Female <input type="radio"/> Other _____
Address			ID / MR#		Gender identification (optional): _____
City	State	Zip Code	Ethnicity <input type="radio"/> Ashkenazi Jewish <input type="radio"/> East Asian <input type="radio"/> African/African American <input type="radio"/> European <input type="radio"/> South Asian <input type="radio"/> Latino <input type="radio"/> Other: _____		
Phone	Email				

Ordering Healthcare Provider			
First Name	Last Name	Title	NPI #
Facility Name			Phone
Facility Address			Fax
City	State	Zip Code	Email
Additional Report Recipients			
Name	Phone	Fax	Email
Name	Phone	Fax	Email

Billing Information		
Insurance Billing		
Insurance Company	Policy #	Group #
Policy Holder First Name	Policy Holder Last Name	Policy Holder DOB
Who is the Policy Holder? <input type="radio"/> Patient <input type="radio"/> Spouse <input type="radio"/> Parent		
Employer's Address		
Institutional Billing	Patient Payment	
An invoice will be sent to the institution listed above. Please contact us for alternate billing.	An invoice will be sent to the patient email provided. Insurance will not be billed.	

Patient Sample Information		
Sample Type	Sample Will Be Collected	Collection date
<input type="radio"/> Saliva <input type="radio"/> Assisted saliva <input type="radio"/> Blood <input type="radio"/> Genomic DNA <input type="radio"/> Other: _____	<input type="radio"/> In-house <input type="radio"/> By Variantyx	____/____/____



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### Form instructions:

- ✓ Review the information on pages 1-4
- ✓ The patient or legal guardian must sign on page 1
- ✓ When submitting comparator samples, the relative(s) must sign on page 5

### Patient Consent

I have discussed the Genomic Unity® test with my healthcare provider including the purpose and procedure, risks, benefits and alternatives. I have been given an opportunity to ask questions about the test, and any questions I had were answered to my satisfaction. I acknowledge that I have sufficient information and understanding to give this informed consent.

1. I give permission to Variantyx and their affiliates to extract and sequence my DNA and perform genetic testing as described.
2. I give permission for my anonymized DNA to be used by Variantyx and their affiliates for test development or improvement, internal validation, orthogonal variant confirmation at an outside referral laboratory and/or quality assurance and training purposes.
3. I give permission for my anonymized sample and clinical information to be included in variant and allele frequency databases and publications. My name or other personal identifying information will not be used in or linked to any databases or publications.
4. In the case of direct insurance billing: I acknowledge that the information provided by me is true and correct. I authorize my healthcare provider and/or insurer to share medical information with Variantyx related to my condition, diagnosis and treatment as relevant to my genetic testing, as well as information about my healthcare plan benefits. I authorize Variantyx to release my medical information concerning my testing to my insurer. I authorize Variantyx to be my designated representative for purposes of appealing any denial of benefits as needed and to request additional medical records for this purpose. I understand that Variantyx will notify me if my out of pocket costs are determined to exceed \$100. I authorize my insurance benefits to be paid directly to Variantyx. I understand that I am responsible for sending Variantyx any and all of the money that I receive directly from my insurer in payment for this test.
5. In the case that independent pre-test and/or post-test genetic counseling is required by my insurance provider and/or physician, I agree, by signing this consent form, to have DNAVisit, a third party, facilitate the genetic counseling services. By signing this consent form, I authorize Variantyx to release my contact information and any medical information necessary to DNAVisit, as well as authorize communication and sharing of information between DNAVisit and my referring physician, in order to complete pre-test and/or post-test genetic counseling. Information about DNAVisit is available at <https://www.dnavisit.com/>.
6. I ☐ give / ☐ do not give permission for Variantyx to contact me or my healthcare provider about research studies. If no option is selected, no contact will be made.
7. Regarding Secondary Findings and Other Incidental Findings (only available with the Genomic Unity® Whole Genome Analysis and Genomic Unity® Exome Analysis tests, and only for the patient):  
I choose ☐ to receive / ☐ not to receive Secondary (ACMG) Findings  
I choose ☐ to receive / ☐ not to receive Other Incidental Findings  
No selection above will default to an opt-out option and findings in these categories will not be returned to you.
8. For NY state residents: ☐ By checking this box I give permission for Variantyx and their affiliates to retain any remaining sample longer than 60 days for testing completion, test development/improvement, internal validation, orthogonal variant confirmation at an outside referral laboratory and/or quality assurance and training purposes.

Patient (or authorized individual) first name

Last name

Patient (or authorized individual) signature

Date





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## Test Information

The benefits and risks of the Genomic Unity® test are explained below. It is recommended that you receive genetic counseling from a licensed healthcare provider who can answer your questions about genetic testing and provide information about alternatives. Information about genetic counselors in your area is available at <https://www.nsgc.org/>.

## Background

The purpose of genetic testing is to identify changes in the DNA sequence that are the cause of an affected individual's condition. This test uses a PCR-free protocol that produces comprehensive and consistent coverage of all exons and non-coding regions in an individual's genome. When applicable to familial samples, whole exome protocols that target exons may be used for comparison to the proband. The resulting data is subjected to *in-silico* analyses optimized for small sequence changes (single nucleotide variants and deletion/insertions), structural variants in chromosomes (deletions, duplications, copy number variants), short tandem repeats (STRs) and mitochondrial variants (single nucleotide variants and small deletion/insertions and large deletions). The Genomic Unity® Whole Genome Analysis test considers mitochondrial variants from the mitochondrial genome as well as most variant types overlapping the exome traditionally reported using other methodologies, excluding technically challenging variants listed in the limitations of testing. The Genomic Unity® Exome Analysis test considers most variant types overlapping the exome traditionally reported using other methodologies, excluding technically challenging variants listed in the limitations of testing. The Genomic Unity® Constitutional Genome-Wide Copy Number Variant Analysis test considers structural variants only and small sequence changes intersecting reportable copy number variants when requested. The Genomic Unity® Mitochondrial Genome Analysis test considers mitochondrial variants from the mitochondrial genome only, and therefore does not include nuclear encoded genes. All other tests consider variants in or overlapping a subset of genes which are described in brief in the Targeted Analyses section of the test requisition form and in more detail on the individual test information web page indicated. When a Custom Analysis is specified, only variants in or overlapping the listed gene(s) specified are considered and only for small sequence changes, deletion/duplications and short tandem repeats as applicable to the gene. All tests are focused on rare variants. When noted for the specified analysis, this test uniquely assesses tandem repeats in genes involved in early-onset intellectual disability (*AFF2*, *DIP2B*, *FMR1*), adult-onset movement disorders with or without cognitive involvement (*AR*, *ATN1*, *ATXN1*, *ATXN2*, *ATXN3*, *ATXN7*, *ATXN8OS*, *ATXN10*, *C9ORF72*, *CACNA1A*, *CNBP*, *CSTB*, *DMPK*, *FMR1*, *FXN*, *NOP56*, *NOTCH2NLC*, *PABPN1*, *PPP2R2B*, *TBP*) and/or other disorders (*PHOX2B*, *TCF4*). Expanded alleles will be reported for these genes when relevant to the patient's clinical symptoms. Based on recommendations by the ACMG, the *JPH3* and *HTT* genes are excluded from this analysis by default, but may be included if a specialized consent form has been signed by the patient/guardian and ordering clinician. Access the form at <https://www.variantyx.com/HTT-JPH3-Consent/>.

The positive predictive value of this test ranges from 0.99676 to 0.99931 depending upon the specific assay selected. Additional information about the Genomic Unity® test is available from your healthcare provider and on the Variantyx website at <https://www.variantyx.com/>. Adult-onset disorders not related to the indication for testing, and therefore representing predictive testing, are not reported with this test. Requests for predictive, carrier and other non-diagnostic genetic testing are available by ordering the Genomic Inform® test.

## Technical Limitations

Genetic testing is accurate, but may not always identify a genetic variant even though one exists. This test attempts to evaluate the entire DNA sequence (within the scope described for the test), but may not be able to detect all DNA changes due to limitations in current technology. Certain regions of the DNA may not be well covered. Certain variant types may not be detectable such as methylation abnormalities, variants in genes with highly homologous pseudogenes and variants in regions that are difficult to assay based on current technology. Unusual circumstances including bone marrow transplantation, blood transfusion, and variants that exist in only a small fraction of cells (mosaicism) may interfere with variant identification. Deletions, duplications and copy number variants between 50 and 300 nucleotides are detected with a lower sensitivity. The false negative rate for repeat expansions has not been determined for the following genes: *AFF2*, *ATXN10*, *CNBP*, *CSTB*, *DIP2B*, *JPH3*, *NOP56*, *NOTCH2NLC*, *PHOX2B*, *TBP*, *TCF4*. The following genes can be assessed for normal repeat ranges only: *DIP2B*, *NOTCH2NLC*, *TCF4*. Repeat counts above 45 will be reported as indeterminate. For dominant repeat expansion disorders parental inheritance will not be reported on the initial report. When exome protocols are applied to familial samples, repeat expansions and most deletions, duplications and copy number variants will not be detected. Any additional test specific limitations are noted on the individual test information web page indicated. Additionally, interpretation of the results is limited by the current medical understanding of disease and available scientific information. This test requires high-quality DNA. In some cases, an additional sample may be needed if the volume, quality and/or condition of the initial sample is not adequate. Samples submitted as genomic DNA will only be processed if the extraction was performed in a CLIA/CAP accredited laboratory. This test does not consider somatic variants.

## Possible Test Results

**Positive result** - A positive result indicates that one or more genetic variants were identified that either explain or partially explain the cause of the disorder or indicate an increased risk of developing the disorder in the future. Individuals with positive results may wish to consider further independent testing and/or consultation with their physician or genetic counselor.

**Negative result** - A negative result indicates that no genetic variant explaining the disorder was identified by this test. This reduces the likelihood of, but does not exclude the possibility of, the disorder being genetic in nature.

**Uncertain result** - A variant of uncertain significance was identified by this test. This means that a genetic variant was identified, but based on available information in the medical literature and research and scientific databases it is not certain whether the variant may cause the disorder. The variant could be a normal genetic difference that does not cause the disorder. Without further information, the effects of the variant cannot be known and an "uncertain/clinically inconclusive" result may be reported. The uncertainty may be resolved over time if additional information



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becomes available. Periodic reanalysis of the sequence data or further analysis, including testing of additional family members, may be recommended.

**Indeterminate result** - An indeterminate result indicates that there were relevant genetic variant(s) identified in the analysis, but that it is uncertain whether they are true variants or artifacts. Furthermore, it is considered that a repeat test will not resolve the technical uncertainty and orthogonal confirmation is necessary to resolve the result.

**Inconclusive result** - A technically inconclusive result indicates that there was an issue with the patient sample that resulted in data that the lab cannot interpret. It is considered that a repeat test will likely resolve the technical uncertainty and therefore a repeat sample is recommended to complete the analysis.

## Reporting Standards

All reportable variants in the clinical report will be categorized as pathogenic, likely pathogenic or a variant of uncertain significance (VUS) utilizing the American College of Medical Genetics and Genomics (ACMG)/Association for Molecular Pathology (AMP) guidelines as published by Richards et al 2015 (for more information see: <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4544753/>). Variants may have a strong phenotypic correlation with the reported patient phenotype(s) and be considered a strong causal candidate for the disorder or may have some phenotypic overlap with the reason for testing but not be considered the sole genetic cause for the phenotype(s) in the patient. Both types of variants may be reported. Even if this test finds DNA changes that are responsible for the reported symptoms, the testing may not completely predict the severity of the disorder, possible future problems, or response to treatment.

## Reporting of Unrelated Findings

Secondary (ACMG) Findings and/or Other Incidental Findings are only available for Genomic Unity® Whole Genome Analysis, Genomic Unity® Exome Plus Analysis and Genomic Unity® Exome Analysis and are not available to relatives, with the exception of the reported parental inheritance of the variants identified in the patient. No specific parental results are issued as a separate report under the family member's name. If the patient chooses to receive secondary or other incidental findings, those findings will be included in a separate section of the clinical report.

With this test related findings are reported, such as genetic findings useful for the current diagnosis of the disease that initially led to the analysis and any clinically relevant genetic findings, which may have immediate benefits for the patient related to present diseases or clinical conditions. However, some unrelated findings may be reported as an option to receive with the report, as listed below, while others such as, pharmacogenomic, high frequency risk alleles, carrier status (heterozygous pathogenic variants in genes associated with autosomal recessive conditions that are not associated with the patient's reported symptoms) and late onset disorders, etc., are outside the scope of testing and would not be typically reported. These different findings and options to receive results are described below.

## Unrelated Findings

Unrelated Findings are findings obtained from genomic sequencing, usually whole genome or exome sequencing, and can be related to conditions that were not the primary reason for testing or findings that can allow one to deduce information as a result of testing that is not directly related to the test. Unrelated findings can be further defined into different types of incidental and secondary findings.

### Unavoidable Incidental Findings (typically reported if present)

Some incidental findings are unavoidable and can be deduced from testing, such as discovering non-paternity when testing the parents of a child in trio analysis or discovering that a parent is a carrier for the condition identified in the child. Other incidental findings are variants in genes that may fit the patient's clinical phenotype but are also related to clinical symptoms unrelated or with a later onset. For example, more than 450 different pathogenic variants have been identified in the *LMNA* gene, which can cause a wide variety of distinct and disparate diseases involving striated muscle (dilated cardiomyopathy, skeletal myopathies), adipose tissue (lipodystrophy syndromes), peripheral nerve (Charcot-Marie-Tooth neuropathy) or multiple systems with accelerated ageing (progerias). These results would likely be reported because they are integral to testing. The possibility of receiving unavoidable incidental findings should be discussed with the patient and family prior to testing, so they are aware that these results, if present, are likely to be returned to them. If the patient does not wish to receive these results, they can decide not to continue with testing.

Patients for whom the Genomic Unity® Whole Genome Analysis, Genomic Unity® Exome Plus Analysis or Genomic Unity® Exome Analysis test is ordered have the choice to opt-in to two additional sets of findings:

### ACMG Secondary Findings

The American College of Medical Genetics and Genomics (ACMG) recommends reporting pathogenic and likely pathogenic variants in a list of genes in both a gene-specific and variant-specific manner. Variantyx evaluates the secondary findings list of genes, the version of which will be listed in the report and can be found on the Variantyx website, [www.variantyx.com/acmg-secondary-findings](http://www.variantyx.com/acmg-secondary-findings). These variants are not typically reviewed during routine processing of patient samples, but are actively sought and reported to the patient. The ACMG recommends reviewing variants in the genes in their recommended list because the genes are related to conditions that are considered 'actionable', meaning that there are steps that can be taken to mitigate the onset or severity of the clinical outcome. It is important to understand that it is possible to have a pathogenic variant but to have it not detected by the assay. In addition, variants of uncertain significance are not reported in these genes. If a variant is of uncertain significance, and later is considered pathogenic, it cannot be determined without a reanalysis of the data.

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### Other Incidental Findings

Other incidental findings are discovered in genes unrelated to the patient's present symptoms, but may have some actionability such as monitoring for possible cardiac implications, increased cancer screening, monitoring of iron levels, have a dietary impact or are diseases for which possible treatment is available (e.g. cardiovascular diseases predisposing to sudden cardiac death). These are genes not on the ACMG list but are similar in that they could impact medical management and decision making.

### Special Consent

To receive results on select genes that are not included in the above, a special consent form is required.

In general the Genomic Unity® tests are for diagnostic purposes and are not offered for predictive testing such as identifying adult onset disorders in an individual who is too young for disease manifestation. In addition, certain genes require additional genetic counseling for the patient and family due to the complexity of the clinical implications and psychological consequences of receiving such results. Therefore, for individuals seeking these results an additional consent is required.

### Testing of Family Samples

In the case of trio and/or larger cohort analysis, and for parental confirmation of singleton analysis, sequencing and analysis of family samples may be used to improve the interpretation of genetic variants identified in the patient's DNA. Variantyx will determine the method (exome or genome) of sequencing used for the familial samples based on the variants identified in the patient's DNA. Accurate interpretation of test results requires accurate assignment of family relationships. Analysis of the sequenced DNA is performed with the assumption that correct family relationships have been provided. Parental samples that fail concordance with the patient (i.e. one parent does not share the expected number of variants with the child) will not be analyzed. Family samples are analyzed only with regard to the patient's condition. Parental inheritance is reported on variants if identifiable, this may include the inheritance of variants related to incidental or secondary findings. However for patients with repeat expansions, parental inheritance may not be reported. Additional counselling for the parents may be recommended prior to reporting parental inheritance of the repeat expansion.

### Patient Confidentiality

To maintain confidentiality, test results will only be released to the ordering healthcare provider or ordering laboratory, and upon your request, to additional healthcare provider(s) indicated on this test requisition form. Test results will only be disclosed to others by your written consent and/or if demanded by a court of competent jurisdiction. It is your responsibility to consider the possible impact of test results on insurance rates, the ability to obtain disability, life or long-term care insurance and employment. The Genetic Information Non-discrimination Act (GINA), enacted by the US Federal Government, provides some protection against discrimination by health insurance companies and employers based on genetic test results, but does not cover life, disability or long-term care insurance. Information about GINA is available at <https://www.genome.gov/10002328>.

Anonymized information obtained from the test may be included in variant and allele frequency databases used to help healthcare providers and scientists understand human disease, as well as in scientific publications. Names and personal identifying information will not be revealed. Separate from the above, if there are opportunities to participate in research relevant to your condition, and you have consented for recontact, Variantyx may contact you or your healthcare provider for research purposes.

### Turnaround Time

The turnaround time (TAT) of this test can be found on the [Variantyx website](#), which begins at the time of sample receipt. For family testing, the timing starts when the last sample is received. Please note that the following scenarios will likely result in extension of the turnaround time (1) when the DNA sample fails QC and is determined to be insufficient for testing, requiring collection of a new sample; (2) when the test is sent for orthogonal confirmation at an external laboratory. In the second scenario, the turnaround time can be expected to be extended by the turn around time of the external laboratory plus 1 week.

### Sample Retention

DNA extracted from submitted samples may be stored for at least 3 months following completion of testing and may be discarded thereafter. Extracted DNA is not returned unless requested prior to testing (additional fees apply). After completion of testing, anonymized DNA may be used for test development and improvement, internal validation, quality assurance and training purposes before being discarded.

NY state residents: No other test shall be performed on this sample except the test ordered by the clinician, unless waived by the patient or authorized individual. In addition, the patient's biological sample will be destroyed within 60 days or upon the completion of testing, unless waived by the patient or authorized individual. Orthogonal confirmation of results at a reference lab cannot be performed unless the patient or authorized individual provides permission to do so.





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### Biological Mother's Information

First Name	Last Name	DOB
Sample Collection Date	Sample Type <input type="radio"/> Blood <input type="radio"/> Saliva <input type="radio"/> Genomic DNA	If affected by the same disorder as the patient please list the clinical symptoms:

### Biological Father's Information

First Name	Last Name	DOB
Sample Collection Date	Sample Type <input type="radio"/> Blood <input type="radio"/> Saliva <input type="radio"/> Genomic DNA	If affected by the same disorder as the patient please list the clinical symptoms:

### Other Relative's Information

First Name	Last Name	DOB
Sample Collection Date	Sample Type <input type="radio"/> Blood <input type="radio"/> Saliva <input type="radio"/> Genomic DNA	If affected by the same disorder as the patient please list the clinical symptoms:
Relationship to Patient Brother:                      Sister:                      Other:		

### Family Member Consent

I have discussed the Genomic Unity® test with my healthcare provider including the purpose and procedure, risks, benefits and alternatives. I have been given an opportunity to ask questions about the test, and any questions I had were answered to my satisfaction. I acknowledge that I have sufficient information and understanding to give this informed consent.

1. I give permission to Variantyx and their affiliates to extract and sequence my/my relative's DNA and perform genetic testing for the purpose of improving the interpretation of genetic variants identified in the patient's DNA.
2. I give permission for my anonymized DNA to be used by Variantyx and their affiliates for test development or improvement, internal validation, orthogonal variant confirmation at an outside referral laboratory and/or quality assurance and training purposes.
3. I give permission for my anonymized sample and clinical information to be included in variant and allele frequency databases and publications. My name or other personal identifying information will not be used in or linked to any databases or publications.
4. For NY state residents: ☐ By checking this box I give permission for Variantyx and their affiliates to retain any remaining sample longer than 60 days for testing completion, test development/improvement, internal validation, orthogonal variant confirmation at an outside referral laboratory and/or quality assurance and training purposes.

Biological mother's first name	Last name
Biological mother's signature	Date
Biological father's first name	Last name
Biological father's signature	Date
Other relative's first name	Last name
Other relative's (or authorized individual) signature	Date